
(12) UK Patent Application (19) GB (11) 2 120 007 A

- (21) Application No **8310253**
(22) Date of filing
 15 Apr 1983
(30) Priority data
(31) **368967**
(32) **16 Apr 1982**
(33) **United States of America**
 (US)
(43) Application published
 23 Nov 1983
(51) **INT CL³ G01N 27/62**
 C07B 23/00
(52) Domestic classification
 H1D 21X
 G1B BC
 U1S 1464 1483 1484
 2159 G1B H1D
(56) Documents cited
 None
(58) Field of search
 H1D
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**(54) Isotope determination by
mass spectrometry**

(57) A mass spectrometry method for determining the absolute value of a given isotopic ratio of an unknown sample and/or the difference of isotope content between an unknown sample and a reference, comprises a) obtaining with or without chemical reaction a substrate capable of providing a fragmentation characteristic of the presence or absence of a specified isotope and usable to retrace the parent ions by the metastable ions technique consisting of accelerating voltage scan, b) introducing said substrate into the source of a mass spectrometer followed by the ionization of said substrate, c) retracing with metastable ions technique the parent ions of a daughter ion resulting from the loss of a neutral frag-

ment characteristic of the presence or absence of the specified isotope in the substrate, d) comparing the relative intensities as expressed by peak areas or heights, of the metastable transitions between, first, said daughter ion and the substrate parent ion, and second, said daughter ion and a transition characteristic of another isotope of well known abundance and usable as internal reference.

For example, a suitable substrate for deuterium determination in water samples is 1-propanol, and for oxygen-18 determination in water samples is ethyl propanoate (propanoic acid, ethyl ester).

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SPECIFICATION

Process for the determination of isotopes by mass spectrometry

5 BACKGROUND OF THE INVENTION

Isotope determination in water or other specific substrate is an important tool in chemistry as well as in biochemistry. Among the most striking applications, I found: the quality control of the foods, the determination of the origin of wines, the geochemical history of organic material, numerous applications in nuclear chemistry and physics, the study of metabolism, and the measure of the total body water living organisms. This latter method, for example, is based on isotope dilution with deuterium or oxygen-18 and is principally used for the understanding of the control of energy balance in humans, for the study of metabolism, or in estimating the body fat content in animals.

Although it is interesting to quantify many isotopic ratios, the most commonly measured are: D/H, carbon 13/12, oxygen 18/16, nitrogen 15/14, and sulphur 34/32.

As an example, several methods are available to quantify the D/H ratio in water: infrared spectrometry, freezing point, falling drop, gas chromatography, NMR spectrometry, and mass spectrometry. These methods have been widely discussed in the literature and only the latter exhibits the required accuracy and sensitivity to be used to detect the small changes in isotopic distribution of deuterium in water at parts-per-million (ppm) level or less.

Two principal approaches can be used by mass spectrometry: first, the direct measurement of the ratio of either m/z 19/18 (HOD/ HOH) or m/z 19/20 (HOD/HO¹⁸H) ion abundances in water samples, provided that the exact amount of oxygen isotopes can be estimated (Anal. Chem. 1953, 25, pp 130-134); second, the measurement of the D/H ratio in gaseous hydrogen samples (Biomed. Mass Spectrom., 1977, 4, pp 82-87).

Both methods suffer some drawbacks. The first is dependent on the accuracy of the evaluation of H₃O⁺ abundance in peak m/z 19 and, although its influence can be minimized by increasing the source repeller voltage to a high value, this evaluation is still dependent on the source pressure, a parameter among the most difficult to control accurately. The second method is by far the most accurate but involves the use of a very sophisticated mass spectrometer especially designed for the measurement of the isotopic content. Such an instrument is expensive and often unavailable. Moreover, it requires a time consuming and tedious decomposition of water into pure gaseous hydrogen, which is usually done with a uranium furnace (Anal. Chem. 1980, 52, pp 2232-2243), and comparison with an external standard which has itself some inherent inaccuracies in its isotope composition due to the industrial method used for hydrogen preparation.

Similar difficulties occur with the other cited isotopic ratios which are usually measured as carbon dioxide for carbon 13/12 and oxygen 18/16, as molecular nitrogen 15/14, and as sulphur dioxide for sulphur 34/32.

Accordingly, it would be highly desirable if an unambiguous method suitable for a standard double focusing mass spectrometer was developed without or with minimum sample transformations and usable for as many as possible isotopic ratios.

Furthermore, it would appear highly desirable if a faster and more general method for mass spectrometry determination of: first, deuterium and oxygen-18 in water, and second, for other isotopes in appropriate substrates, can be developed for use with standard double focusing mass spectrometer at ppm level from natural abundance up to high concentration of the studied isotopes. It would also appear to be highly desirable if a method could be developed which would be devoid of the prior art methods while also avoiding the necessity of excessive purification of the sample to be tested.

SUMMARY OF THE INVENTION

In accordance with the present invention, there is provided a novel method, based on a concept developed on deuterium and oxygen-18 content determination in water samples, for determining by mass spectrometry the isotope content in an appropriate substrate.

More specifically, the novel method of the present invention comprises:

- a) either introducing the studied isotope as a single entity (e.g. H \rightleftharpoons D) or as a function (e.g. O¹⁶H = O¹⁸H) with an appropriate substrate, or using a chemical reaction to yield an appropriate substrate involving the studied isotope,
- b) providing a molecular fragmentation characteristic of the presence or absence of the studied isotope and usable to quantify the said isotope content through the use of either the metastable ions technique (accelerating voltage scan) or another appropriate method to retrace parent ions.

The mixture is then introduced into the source of a mass spectrometer and then ionized. The transitions between the daughter ion resulting from the loss of a neutral fragment characteristic of the presence or absence of the said isotope from the substrate and the parent ions are then

retrace d.

Finally, the relative intensities of the specified transitions as measured by the ratio of peak areas or heights are used, with or without calibration with standards, to determine the difference of isotope content between the unknown sample and a reference.

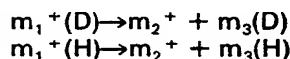
DETAILED DESCRIPTION OF THE INVENTION

The method of the present invention takes advantage of the capability of either double focusing mass spectrometers to retrace the parent ions of a given daughter ion by scanning up the accelerating voltage, the magnetic and electrostatic sectors being set on the daughter ion parameters, or multiple stages mass spectrometer to retrace the parent ions of a given daughter ion by scanning the first stage, the second stage being set on the daughter ion parameters. The main advantage of such a method is its extreme specificity and thus its ability to circumvent all the inherent problems of the isotope interferences and the presence of other concurrent ionic species. Moreover, because of the specificity of the approach, it allows one to simplify to a great extent the preparation of the sample and does not necessarily involve an especially designed and dedicated mass spectrometer for isotope measurements. Finally, the developed concept being in itself exceptionally stable, it allows to reduce to a great extent the number of sophisticated and expensive accessories necessitated by the conventional methods. This is due to the fact that the present invention is based on only one specific ionic species instead of at least two different kind of ions for the conventional method.

The method can be best described on two specific examples.

A. Deuterium determination in water samples.

For a given m_2^+ daughter ion the fragmentation will be:

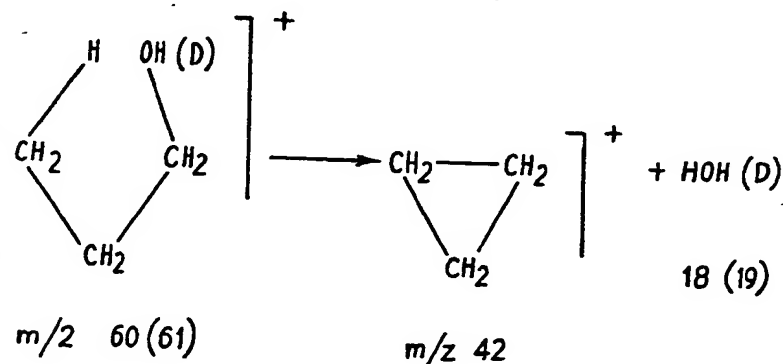


with $m_1^+(D) = m_1^+(H) + 1$ and $m_3(D) = m_3(H) + 1$ and where parent ions are $m_1^+(D)$ or $m_1^+(H)$ depending on the presence of deuterium or hydrogen atom, respectively. The only limiting factor for an accurate measurement is the necessity to have to intense metastable transitions to link together daughter and parent ions.

It will be appreciated that the direct measurement of $18^+(H) \rightarrow 17^+$ and $19^+(D) \rightarrow 17^+$ transitions in water is not easily accessible mainly because of the weakness of the corresponding metastable peaks.

Accordingly, in accordance with the present invention there must be used a substrate which is suitable for the interchanging reaction $RH + DOH \rightleftharpoons RD + HOH$ and the measure of the RD/RH ratio after equilibration. The substrate could be an oxygen containing substrate or another organic capable of interchanging hydrogen and deuterium in an aqueous medium and capable of providing a fragmentation characteristic of the presence or absence of an interchanged deuterium so as to retrace the parent ions by either the metastable ions technique or another appropriate method.

As an example of a suitable substrate there may be mentioned 1-propanol. As a primary alcohol with three carbon atoms it undergoes the following fragmentation:



The parent ions are m/z 60 or 61 depending on the very isotope nature of the hydroxylic hydrogen. Thus, providing that, first, the isotope effect during the interchanging reaction with 1-propanol can be kept constant, second, the deuterium content of the terminal methyl is constant, and third, the isotope effect during fragmentation is negligible or constant, measure-

ment of the intensity ratio between $42^+ \rightarrow 61^+$ and $42^+ \rightarrow 60^+$ transitions (ROD/ROH ratio) is a direct measure of the deuterium content of the hydroxyl group of 1-propanol.

The possible isotope effect during the deuterium interchange reaction between water and 1-propanol can be circumvented by an appropriate control of the inlet system temperature. This is not a limiting factor because, first, the interchange reaction is extremely fast even at room temperature, and second, the septum inlet used for sample injection can be easily kept at an accurate temperature thus keeping this effect, if it exists, constant.

Natural deuterium and oxygen-18 abundances being 0.015% and 0.04%, respectively, one can expect from the above fragmentation of pure 1-propanol a basic ratio ROD/ROH of $2 \times 0.015 + 0.04 = 0.070$ for the studied transition. It has been found ROD/ROH = 0.0634 ± 0.00005 ($n = 10$, each of them being the sum of 5 different readings), the difference from the expected value is attributed to the isotopic effect. Standard error is taken as $s/n^{1/2}$, where s is the standard deviation and n the number of samples. The validity of the method is confirmed by an additional piece of evidence: pure 1-propanol allowed to interchange its hydroxylic hydrogen with an equal volume of deuterium depleted water (natural abundance divided by 100) exhibits a ROD/ROH ratio equal to 0.0465 ± 0.0006 ($n = 9$).

Calibration curve with standards from 0.025% to 98.85% deuterium above natural level exhibits their best fit for a polynomial regression of degree 2 ($R^2 = 0.9994$). Calibration curves for deuterium concentrations ranging from 0% (pure water) to 0.1% can be reduced to a polynomial regression curve of degree 1 ($R^2 = 0.99988$). Table I shows the reproducibility of the measure. Table II shows the stability of the measure with a series of standards ranging from 0.0471% absolute, as exhibited when using deuterium depleted water, to 0.147% deuterium. Higher concentration of deuterium (greater than 0.1%) were measured by comparison of peak heights and areas of the $42^+ \rightarrow 60^+$ and $42^+ \rightarrow 61^+$ transitions of 1-propanol while low deuterium concentrations (less or equal to 0.1%) were measured by comparison of $42^+ \rightarrow 61^+$ and $42^+ \rightarrow 62^+$ transitions. The latter transition, used as an internal reference for low deuterium content, is based on the assumption that the water molecule lost during the specified transition can involve an oxygen-18 atom. This assumption holds true because oxygen-18 has a well known natural abundance of 0.204%. Thus, it can serve as an internal reference for an absolute deuterium determination in the specified conditions. Water standards were allowed for a few seconds to interchange deuterium and hydrogen with an equivalent volume of pure 1-propanol before injection into the source of the mass spectrometer.

Calibration curve with standards from 0.025% to 98.85% deuterium above natural level exhibits their best fit for a polynomial regression of degree 2 ($R^2 = 0.9994$). Calibration curves for deuterium concentrations ranging from 0% (pure water) to 0.1% can be reduced to a polynomial regression curve of degree 1 ($R^2 = 0.99988$). Table I shows the reproducibility of the measure.

TABLE I
REPRODUCIBILITY

	Reference %	Sample %
45	0.06013	0.06164
	0.06032	0.06170
	0.06022	0.06152
mean	0.06022	0.06162
S.D.*	0.00010	0.00027
50 ppm	0.0	16.0

*S.D.: Standard deviation

Pure 1-propanol was taken as reference. The unknown sample is a wine "St. Emilion appellation contrôlée". The deuterium content above natural level was measured by comparison of transitions $42^+ \rightarrow 61^+$ and $\rightarrow 62^+$ with oxygen-18 natural abundance (0.204%) as internal standard.

Table II shows the stability of the measure with a series of standards ranging from 0.0471% absolute, as exhibited when using deuterium depleted water, to 0.14% deuterium.

TABLE II
STABILITY OF THE MEASURE

5	Deuterium %	S.D.*	Error %	n	5
	0.0471 (depleted)	0.0002	0.4	4	
10	0.0590 (pure)	0.0001	0.2	5	10
	0.0641	0.0002	0.4	5	
	0.0687	0.0002	0.2	5	
	0.0813	0.0003	0.4	5	
15	0.1043	0.0002	0.2	5	15
	0.1471	0.0003	0.2	5	

*S.D.: Standard deviation

20 All the standards used were made by dilution from a stock solution of 0.1471% deuterium (about 1000 ppm above natural level). Deuterium depleted water was purchased from Sigma (natural level $\times 100$). n is the number of lectures for a given sample. 20

Higher concentration of deuterium (greater than 0.1%) were measured by comparison of peak heights and areas of the $42^+ \rightarrow 60^+$ and $42^+ \rightarrow 61^+$ transitions of 1-propanol while low deuterium concentrations (less or equal to 0.1%) were measured by comparison of $42^+ \rightarrow 61^+$ and $42^+ \rightarrow 62^+$ transitions. The latter transition, used as internal reference for low deuterium content, is based on the assumption that the water molecule lost during the specified transition can involve an oxygen-18 atom. This assumption holds true because oxygen-18 has a well known natural abundance of 0.204%. Thus, it can serve as an internal reference for an absolute deuterium determination in the specified conditions. Water standards were allowed for a few seconds to interchange deuterium and hydrogen with an equivalent volume of pure 1-propanol before injection into the source of the mass spectrometer. 25 30

B. Oxygen-18 determination in water sample

A similar approach has been developed for oxygen-18 determination from water samples. 35
Oxygen from water samples is selectively introduced into the carbonyl group of ethyl propanoate (propanoic acid, ethyl ester) through the hydrolysis of its triethyl ortho propionate ester derivative under acidic condition at room temperature. The reaction is highly reproducible and usable on a routine basis. The obtained ethyl propanoate is then injected into the instrument ion source through the septum inlet. Ethyl propanoate exhibits an intense transition 40 resulting from the loss of water from the molecular ion and involving selectively the carbonylic oxygen. Thus, depending on the presence or the absence of oxygen-18, the studied transition will be $84^+ \rightarrow 102^+$ or $84^+ \rightarrow 104^+$, respectively. Then, the transition intensities are used as described above for deuterium determination. 40

The calibration curve with standards from 0 to 1.5% oxygen-18 above natural level exhibits 45 their best fit for a polynomial regression of degree 2 ($R^2 = 0.9997$). Calibration curves of oxygen-18 concentration ranging from 0% (pure water) to 0.5% can be reduced to a polynomial regression curve of degree 1 ($R^2 = 0.99997$). Table III shows the reproducibility of the measure.

TABLE III

50 OXYGEN-18 REPRODUCIBILITY

	Reference %	Sample %	
55	0.2003	0.2244	55
	0.2001	0.2250	
	0.2004	0.2246	
	0.2004	0.2236	
	0.2007	0.2246	
60 mean	0.2004	0.2244	60
S.D.*	0.0002	0.0005	

*S.D.: Standard deviation

65 Tap water was taken as reference. The unknown sample is a standard involving 0.050 65

percent oxygen-18. The oxygen-18 content was measured by comparison of transitions $84^+ \rightarrow 102^+$ and $84^+ \rightarrow 104^+$.

Table IV shows the stability of the measure with a series of standards ranging from 0% as exhibited when using tap water to 1.5% oxygen-18.

5

TABLE IV
STABILITY OF THE MEASURE

Oxygen-18	S.D.	Error
%		%
0.1875	0.0004	0.10
0.1905	0.0002	0.05
0.1941	0.0001	0.02
0.2244	0.0002	0.04
0.2622	0.0006	0.10
0.5504	0.0031	0.25
0.8852	0.0013	0.07
1.0960	0.0030	0.12

20

Oxygen-18 standards are expressed as the ratio of their transitions intensities $84^+ \rightarrow 104^+$ vs $84^+ \rightarrow 102^+$.

The percentage of error is expressed as $S.D./\sqrt{n} \times 100$ with $n = 5$.

25 All the standards used were made by dilution from a stock solution of 1.5% oxygen-18 above natural level. 25

It should be emphasized that the accuracy of the present invention is substantially of the same order of magnitude as that of the more time-consuming and sophisticated conventional methods. Yet, the results obtained with the present invention do not require any specific accessories and any data acquisition system. Moreover, the time required for one experiment was reduced to seconds instead of up to 15 minutes for most of the conventional methods. 30

C. The same procedure can be applied for other isotope determination providing that an appropriate substrate is used for retracing parent ions resulting from the loss of a neutral fragment involving the studied isotope. It should be emphasized that the accuracy of the present invention is substantially of the same order of magnitude as that of the more time consuming and sophisticated conventional method. Yet, the results obtained with the present invention do not require any specific accessories and any data acquisition system. Despite that, the time required for one experiment was reduced to seconds instead of fifteen minutes for most of the conventional methods. 35

40 CLAIMS 40

1. A mass spectrometry method for determining the absolute value of a given isotopic ratio of an unknown sample and/or the difference of isotope content between an unknown sample and a reference, which comprises

45 a) obtaining with or without chemical reaction a substrate capable of providing a fragmentation characteristic of the presence or absence of a specified isotope and usable to retrace the parent ions by either the metastable ions technique consisting of the accelerating scan, or another technique using multiple stages mass spectrometers, 45

b) introducing said substrate into the source of a mass spectrometer followed by the ionization of said substrate, 50

c) retracing with metastable ions technique or other technique the parent ions of a daughter ion resulting from the loss of a neutral fragment characteristic of the presence or absence of the specified isotope in the substrate, 50

d) comparing the relative intensities, as expressed by peak areas or heights of the transitions between, first, said daughter ion and the substrate parent ion, and second, said daughter ion and a transition characteristic of another isotope of well known abundance and usable as internal reference. 55

2. A mass spectrometry method for determining the absolute value of deuterium content of an unknown aqueous sample and/or the difference of deuterium content between an unknown aqueous sample and a reference, which comprises 60

a) either mixing a given volume of an aqueous sample with a substrate capable of interchanging hydrogen and deuterium in an aqueous medium, or trapping hydrogen and deuterium from water samples by chemical reaction, and capable of providing a fragmentation characteristic of the presence or absence of a specific deuterium and usable to retrace the parent ions by either the metastable ions technique consisting of the accelerating voltage scan, or other 65

technique using multiple stages mass spectrometers,

b) introducing said mixture into the source of a mass spectrometer followed by the ionization of said mixture,

5 c) retracing with either the metastable ions technique or another appropriate technique the parent ions of a daughter ion resulting from the loss of a neutral fragment characteristic of the presence or absence of a specific deuterium in the substrate, 5

d) comparing the relative intensities, as expressed by peak areas or heights, of the transitions between, first, said daughter ion and the substrate parent ion, and second, said daughter ion and the substrate parent ion plus one mass unit, or the latter transition and a transition 10 characteristic of another isotope of well known abundance and usable as internal reference. 10

3. A method according to Claim 1 or 2, wherein the substrate is 1-propanol.

4. A mass spectrometry method for determining the absolute value of oxygen-18 content of an unknown aqueous sample and/or the difference of oxygen-18 content between an unknown aqueous sample and a reference, which comprises

15 a) introducing oxygen-18 from water samples into the carbonyl group of ethyl propanoate by hydrolysis of its triethyl ortho propionate ester derivative, and capable of providing a fragmentation characteristic of the presence or absence of a specific oxygen-18 and usable to retrace the parent ions by either the metastable ions technique consisting of the accelerating voltage scan, or other technique using multiple stages mass spectrometers, 15

20 b) introducing said mixture into the source of a mass spectrometer followed by the ionization of said mixture, 20

c) retracing with either the metastable ions technique or another appropriate technique the parent ions of a daughter ion resulting from the loss of a neutral fragment characteristic of the presence or the absence of a specific oxygen-18 in the substrate,

25 d) comparing the relative intensities, as expressed by peak areas or heights, of the transitions between, first, said daughter ion and the substrate parent ion, and second, said daughter ion and the substrate parent ion plus two mass units, or the latter transition and a transition characteristic of another isotope of well known abundance and usable as internal reference. 25

5. A method according to Claim 1 and 4, wherein the substrate is ethyl propanoate 30 (propanoic acid, ethyl ester). 30

Printed for Her Majesty's Stationery Office by Burgess & Son (Abingdon) Ltd.—1983.

Published at The Patent Office, 25 Southampton Buildings, London, WC2A 1AY, from which copies may be obtained.

